(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication:

27.02.2002 Bulletin 2002/09

(51) Int Cl.7: C07C 29/151

(21) Application number: 01119645.8

(22) Date of filing: 21.08.2001

(84) Designated Contracting States:

AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE TR

Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 21.08.2000 JP 2000249670

(71) Applicant: Mitsubishi Heavy Industries, Ltd. Tokyo (JP) (72) Inventors:

Kobayashi, Kazuto
 2-5-1, Marunouchi, Chiyoda-ku, Tokyo (JP)

lijima, Masaki
 2-5-1, Marunouchi, Chlyoda-ku, Tokyo (JP)

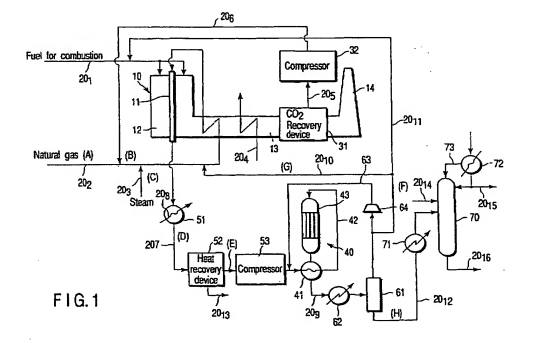
Morita, Kazuhiro
 2-5-1, Marunouchi, Chiyoda-ku, Tokyo (JP)

(74) Representative: HOFFMANN - EITLE
Patent- und Rechtsanwälte Arabellastrasse 4
81925 München (DE)

(54) Method of manufacturing methanol

(57) A natural gas having steam and carbon dioxide added thereto is supplied into a reaction tube (11) of a reformer (10) provided with the reaction tube (11) and a combustion radiation section (12) for heating the reaction tube by combustion of a fuel so as to carry out a reaction between the hydrocarbon contained in the natural gas and the steam added to the natural gas, thereby forming a synthetic gas containing hydrogen, carbon

monoxide and carbon dioxide. Then, the synthetic gas is subjected to a reaction in the presence of a methanol synthesizing catalyst. A crude methanol obtained is subjected to a gas-liquid separation. Further, a liquid crude methanol separated in the separation is distilled. In the particular methanol manufacturing process, a part of the purge gas obtained in the separation is recycled into the natural gas having steam and carbon dioxide added thereto in advance.



Description

15

20

25

[0001] The present invention relates to a method of manufacturing methanol.

[0002] The conventional method of manufacturing methanol (CH₃OH) is as follows:

(Synthetic Gas Forming Process)

[0003] Natural gas having steam and carbon dioxide added thereto is supplied into a reaction tube of a reformer, the reformer being provided with the reaction tube loaded with, for example, a nickel-based catalyst, and a combustion radiation section for heating the reaction tube by the combustion of a fuel. In this case, the hydrocarbon in the natural gas reacts with the steam so as to generate a synthetic gas containing as main components hydrogen, carbon monoxide and carbon dioxide.

[0004] Since the steam reforming reaction described above is an endothermic reaction, the fuel is combusted in the fuel combustion section so as to heat the reaction tube.

(Crude Methanol Synthetic Process)

[0005] The synthetic gas contained hydrogen, carbon monoxide and carbon deoxide is reacted in the presence of a methanol synthetic catalyst under a predetermined pressure and a predetermined temperature so as to synthesize a crude methanol.

(Recovery Process of Liquid Crude Methanol)

[0006] The crude methanol is subjected to a gas-liquid separation for separation into a liquid crude methanol and a purge gas mainly containing hydrogen. The separated purge gas is recycled into the combustion radiation section so as to be utilized as a part of the fuel.

(Distillation Process)

30 [0007] The liquid crude methanol recovered in the recovery process is distilled in one or a plurality of distillation towers so as to separate the crude methanol into a refined methanol and a waste water containing an organic compound having a boiling point lower than that of methanol (hereinafter referred to as a "low boiling point organic compound"), an organic acid, and an organic compound having a boiling point higher than that of methanol (hereinafter referred to as a "high boiling point organic compound").

[0008] Methanol is manufactured through the processes described above.

[0009] An object of the present invention is to provide a method of manufacturing methanol, in which a purge gas obtained by a gas-liquid separation of a crude methanol and mainly containing hydrogen is recycled to a natural gas having steam and carbon dioxide added thereto so as to utilize the purge gas as a part of the fuel, thereby increasing the methanol yield.

40 [0010] According to a first aspect of the present invention, there is provided a method of manufacturing methanol, comprising:

a synthetic gas forming step, in which a natural gas having steam and carbon dioxide added thereto is supplied into a reaction tube of a reformer provided with the reaction tube and a combustion radiation section for heating the reaction tube by the combustion of a fuel so as to bring about a reaction between the hydrocarbon contained in the natural gas and the steam, thereby forming a synthetic gas containing mainly hydrogen, carbon monoxide and carbon dioxide;

a crude methanol synthetic step for performing the reaction of the synthetic gas in the presence of a methanol synthetic catalyst so as to synthesize a crude methanol;

a gas-liquid separation step for separating the crude methanol into a liquid crude methanol and a purge gas mainly containing hydrogen; and

a distillation step for distilling the liquid crude methanol so as to separate a refined methanol;

wherein a part of the purge gas is recycled into the natural gas having steam and carbon dioxide added thereto in advance.

55

45

50

[0011] In the method of manufacturing methanol according to the present invention, it is desirable for a part or all of the carbon dioxide gas added to the natural gas to be the carbon dioxide recovered from at least one of the combustion waste gas generated from the combustion radiation section of the reformer and the combustion waste gas generated

from the boiler for the steam generation.

10

20

25

30

40

45

50

[0012] In the method of manufacturing methanol according to the present invention, it is acceptable for a part of the purge gas to be recycled into the combustion radiation section of the reformer so as to be utilized as a part of the fuel.

[0013] This summary of the invention does not necessarily describe all necessary features so that the invention may also be a sub-combination of these described features.

[0014] The invention can be more fully understood from the following detailed description when taken in conjunction with the accompanying drawings, in which:

FIG. 1 is a diagram schematically exemplifying a methanol manufacturing plant according to one embodiment of the present invention; and

FIG. 2 is a diagram schematically exemplifying a methanol manufacturing plant for the comparative case.

[0015] The method of the present invention for manufacturing methanol will now be described in detail with reference to the accompanying drawings.

[0016] FIG. 1 is a diagram schematically showing the gist portion of a plant used for the method of the present invention for manufacturing methanol.

[0017] As shown in the drawing, the methanol manufacturing plant comprises a reformer 10 including a steam reforming reaction tube 11, a combustion radiation section 12 arranged to surround the reaction tube 11 for heating the reaction tube 11 by combusting a fuel, and a smokestack 14 communicating with the combustion radiation section 12 through a convection section (waste heat recovery section) 13. A catalyst, e.g., a nickel-based catalyst, is loaded in the reaction tube 11. A fuel introducing fluid passageway 20₁ is connected to the combustion radiation section 12 of the reformer 10.

[0018] A raw material gas introducing fluid passageway 20₂ is connected to the upper end of the reaction tube 11 through the convection section 13 of the reformer 10. It is possible to mount a desufurizer (not shown) to the fluid passageway 20₂. A steam introducing fluid passageway 20₃ is connected to the raw material gas introducing fluid passageway 20₂ upstream of the convection section 13. Further, a fluid passageway 20₄ for circulating, for example, a boiler water extends to cross the convection section 13 of the reformer 10 such that a heat exchange is performed between the combustion waste gas of the convection section 13 and the boiler water circulated within the fluid passageway 20₄ so as to cool the combustion waste gas and, at the same time, to heat the boiler water itself, thereby generating a high pressure steam.

[0019] A carbon dioxide recovery device 31 is arranged in the convection section 13 of the reformer 10 so as to recover carbon dioxide in the combustion waste gas generated from the convection section 13. The carbon dioxide recovery device 31 is connected to a compressor 32 via a fluid passageway 20₅. Further, the compressor 32 is connected to the raw material gas introducing fluid passageway 20₂ upstream of the reformer 10.

[0020] A methanol synthesizing reaction apparatus 40 is arranged downstream of the reformer 10 and is connected to the reaction tube 11 of the reformer 10 via a fluid passageway 20₇. The reaction apparatus 40 comprises a preheater 41 and a methanol synthesizing reactor 43 to which is supplied the synthesized gas from the pre-heater 41 through a circulating fluid passageway 42. A methanol synthesizing catalyst is loaded in the reactor 43. A heat exchanger 51, a heat recovery device 52 and a compressor 53 are mounted to the fluid passageway 20₇ interposed between the reformer 10 and the pre-heater 41 in the order mentioned as viewed from the reformer 10. A fluid passageway 20₈ crosses the heat exchanger 51 so as to heat, for example, the boiler water circulated through the fluid passageway 20₈, thereby generating a high pressure steam.

[0021] The reactor 43-of the methanol synthesizing apparatus 40 is connected to a gas-liquid separator 61 through a fluid passageway 20₉. The pre-heater 41 and a cooler 62 are mounted to the fluid passageway 20₉. The gas-liquid separator 61 is connected to the fluid passageway 20₇ upstream of the pre-heater 41 through a gas circulating fluid passageway 63. A gas compressor 64 is mounted to the gas circulating fluid passageway 63. A purge gas fluid passageway 20₁₀ is branched from that portion of the gas circulating fluid passageway 63 which is interposed between the gas-liquid separator 61 and the gas compressor 64 so as to be connected to the raw material gas introducing fluid passageway 20₂. Also, the purge gas fluid passageway 20₁₀ is branched to form a branched purge gas fluid passageway 20₁₁, which is connected to the fuel introducing fluid passageway 20₁.

[0022] The gas-liquid separator 61 is connected to a distillation tower 70 through a fluid passageway 20₁₂. A liquid crude methanol pre-heater 71 is mounted to the fluid passageway 20₁₂. Further, a condenser 72 is connected to the top region of the distillation tower 70 through a circulating passageway 73.

[0023] How to manufacture methanol will now be described with reference to the manufacturing plant shown in FIG. 1.

(1) Synthesizing Gas Forming Step

[0024] In the first step, a fuel for combustion is supplied into the combustion radiation section 12 of the reformer 10

through the fuel introducing fluid passageway 20₁ for the combustion in the presence of the air so as to heat the inside of the reaction tube 11 to a sufficiently high temperature, e.g., 850 to 900°C. The reaction tube 11 is heated because the reforming reaction carried out within the reformer 10 is an endothermic reaction. The combustion waste gas containing carbon dioxide generated in the combustion radiation section 12 flows through the convection section 13 to reach the smokestack 14. While passing through the convection section 13, the combustion waste gas carries out a heat exchange with a natural gas having steam added thereto and circulated within the raw material gas introducing fluid passageway 20₂ and with the boiler water circulated through the fluid passageway 20₄. Carbon dioxide contained in the cooled combustion waste gas is recovered in the carbon dioxide recovery device 31 so as to be supplied into the compressor 32 through the fluid passageway 20₅. The combustion waste gas having carbon dioxide removed therefrom is discharged to the air atmosphere through the smokestack 14.

[0025] The natural gas containing methane as a main component is supplied into the raw material gas introducing fluid passageway 20₂. In this case, a predetermined amount of carbon dioxide compressed by the compressor 32 is added to the natural gas through the fluid passageway 20₆. Also, a predetermined amount of steam is added to the natural gas, to which carbon dioxide has been added, through the steam introducing fluid passageway 20₃. It is possible to utilize the steam generated by the heat exchange between the boiler water and the synthetic gas, which is carried out in the heat exchanger 51, and by the heat exchange between the boiler water and the combustion waste gas, which is carried out in the convection section 13 of the reformer 10. Further, a predetermined amount of the unreacted gas referred to herein later, which mainly contains hydrogen, is added as a purge gas to the natural gas having carbon dioxide and steam added thereto.

[0026] In adding carbon dioxide and steam to the natural gas, it is desirable for the molar ratio of methane (CH₄) contained in the natural gas to steam (H₂O) to fall within a range of between 1:1.5 and 1:5, and for the molar ratio of methane contained in the natural gas to carbon dioxide (CO₂) to fall within a range of between 1:1 and 1:3.

[0027] The natural gas having carbon dioxide and steam added thereto is circulated through the raw material gas introducing fluid passageway 20₂ and is preheated when passing through the convection section 13 of the reformer 10. Further, the natural gas is heated to a sufficiently high temperature and, then, introduced into the reaction tube 11. [0028] The natural gas containing methane (CH₄) as a main component and having steam and carbon dioxide added thereto, which is supplied into the reaction tube 11 of the reformer 10, is subjected to a steam reformation within the reaction tube 11 in the presence of a catalyst. As a result, methane is mainly subjected to a steam reformation so as to generate a synthetic gas containing hydrogen, carbon monoxide and carbon dioxide, as denoted by reaction formulas (1) and (2) given below:

$$CH_4 + H_2O \Leftrightarrow CO + 3H_2 \tag{1}$$

$$CO + H_2O \Leftrightarrow CO_2 + H_2 \tag{2}$$

[0029] As is apparent from reaction formulas (1) and (2) given above, 4 mols of hydrogen and 1 mol of carbon dioxide are generated by the reaction between 1 mol of methane and 2 mols of steam. In the actual reaction system, however, obtained is a composition close to the chemical reaction equilibrium composition determined by the temperature and pressure at the outlet of the reaction tube 11.

(2) Crude Methanol Synthesizing Step

20

30

45

50

[0030] The synthetic gas formed in the reformer 10 is supplied into the heat exchanger 51 through the fluid passageway 207. In the heat exchanger 51, the boiler water passing through the fluid passageway 208 is heated so as to generate a high pressure steam. At the same time, the synthetic gas itself is cooled and, then, supplied to the heat recovery device 52 so as to be cooled to room temperature. In this case, the steam contained in the synthetic gas is condensed, and the condensed water is taken out through a fluid passageway 2013 so as to be utilized as, for example, a process water.

[0031] The synthetic gas having the condensed water separated therefrom is supplied to the compressor 53 through the fluid passageway 20₇ so as to be compressed to have a pressure adapted for the methanol synthesizing reaction, e.g., 50 to 150 atms. The compressed synthetic gas is further supplied to the pre-heater 41 of the reaction apparatus 40 for synthesizing methanol through the fluid passageway 20₇. In the pre-heater 41, the compressed synthetic gas is preheated to a temperature adapted for the methanol synthesizing reaction, e.g., 200 to 300°C, and, then, further supplied into the reactor 43 loaded with a methanol synthesizing catalyst through the circulating fluid passageway 42. Incidentally, the unreacted gas, which is separated in the gas-líquid separator 61 referred to herein later, is supplied

to the fluid passageway 20₇ upstream of the pre-heater 41 through the gas circulating fluid passageway 63 so as to be mixed with the synthetic gas. Reactions (3) and (4) given below are carried out in the reactor 43 so as to form a composition containing the synthesized methanol:

$$CO + 2H_2 \Leftrightarrow CH_3OH$$
 (3)

$$CO_2 + 3H_2 \Leftrightarrow CH_3OH + 2H_2O \tag{4}$$

[0032] Impurities such as dimethyl ether and ethanol are also formed by side reactions. These impurities are also contained in the liquid crude methanol together with methanol formed by the reactions given above.

[0033] A catalyst such as a copper-based catalyst is used as the methanol synthesizing catalyst. Particularly, it is desirable for the methanol synthesizing catalyst to consist of oxides containing Cu, Zn, Al, Ga and M, M being at least one element selected from the group consisting of alkaline earth metal elements and rare earth elements, the oxides having a high durability under an atmosphere containing a high concentration of carbon dioxide, and the atomic ratio of Cu: Zn: Al: M being 100: 10 to 200: 1 to 20: 0.1 to 20.

(Recovery Step of Liquid Crude Methanol)

[0034] The reaction mixture formed in the reactor 43 is supplied into the pre-heater 41 and, then, into the cooler 62 successively through the fluid passageway 20₉ so as to be cooled to substantially room temperature. In this case, methanol and water contained in the reaction mixture are condensed substantially completely so as to form a liquid crude methanol flowing into the gas-liquid separator 61. In the gas-liquid separator 61, the reaction mixture is separated into the liquid crude methanol and the unreacted gas mainly containing hydrogen.

[0035] The unreacted gas is transferred into the gas compressor 64 through the gas circulating fluid passageway 63. After being compressed in the gas compressor 64, the unreacted gas is circulated into the fluid passageway 20₇ upstream of the pre-heater 41 through the gas circulating fluid passageway 63 so as to be supplied into the reactor 43 together with the synthetic gas. A part of the unreacted gas is supplied as a purge gas into the raw material gas introducing fluid passageway 20₂ through the purge gas fluid passageway 20₁₀ so as to be utilized as a part of the raw materials. Also, the remaining unreacted gas is supplied as a purge gas into the fuel introducing fluid passageway 20₁ through the branched purge gas fluid passageway 20₁₁ so as to be utilized as a part of the fuel in the combustion radiation section of the reformer 10.

(Distillation Process)

5

10

20

25

30

35

40

45

50

[0036] The crude methanol separated in the gas-liquid separator 61 is supplied to the distillation tower 70 through the crude methanol pre-heater 71 mounted to the fluid passageway 20₁₂. Also, a small amount of water is supplied to the distillation tower 70 through the fluid passageway 20₁₄. In the distillation tower 70, the liquid crude methanol is separated under the function of the condenser 72 into a refined methanol having a high purity and a waste water containing the low boiling point organic compound and the high boiling point organic compound formed as by-products. The refined methanol is withdrawn as a product to the outside through a fluid passageway 20₁₅. On the other hand, the waste water is discharged to the outside through a fluid passageway 20₁₆.

[0037] Incidentally, carbon dioxide added to the natural gas is not limited to that recovered from the combustion waste gas formed in the combustion radiation section. It is also possible to utilize carbon dioxide gas recovered from the combustion waste gas generated in, for example, the boiler and discarded in, for example, another factory. In other words, carbon dioxide generated from, for example, other factories, which was discarded in the past, can be effectively utilized as a raw material in the method of the present invention for manufacturing methanol so as to decrease the amount of carbon dioxide discharged to the air atmosphere. In other words, the method of the present invention is effective for suppressing global warming, which is a serious environmental problem nowadays.

[0038] A single distillation tower is used in the embodiment described above. However, it is possible to use a plurality of distillation towers.

[0039] As described above, in the method of the present invention, the purge gas mainly containing hydrogen, which is obtained by the gas-liquid separation of the crude methanol, is partly recycled into the natural gas having steam and carbon dioxide added thereto so as to utilize the purge gas as a part of the raw material. In the conventional method, however, the purge gas is recycled into the combustion radiation section of the reformer so as to utilize the purge gas as a part of the fuel. It follows that the method of the present invention permits forming a synthetic gas containing

higher molar ratios of CO and H_2 in the forming process of the synthetic gas, compared with the conventional method. It follows that the method of the present invention permits increasing the crude methanol production and the refined methanol production.

[0040] It should also be noted that carbon dioxide recovered from the combustion waste gas generated from the combustion radiation section of the reformer (or the combustion waste gas generated from the boiler) can be utilized in the present invention so as to decrease the amount of carbon dioxide discharged to the outside in the methanol manufacturing process. It follows that the methanol manufacturing plant can be made economical in the case where the tax on the carbon dioxide gas discharge is introduced and where the carbon dioxide gas discharge is regulated.

[0041] A preferred embodiment of the present invention will now be described with reference to the methanol manufacturing plant shown in FIG. 1.

(Example 1)

20

30

35

40

[0042] A fuel, e.g., a natural gas, was introduced into the combustion radiation section 12 of the reformer 10 at a flow rate of 540 kmoVhr so as to be combusted in the combustion radiation section 12 in the presence of the air. Also, a natural gas, steam and carbon dioxide recovered from the combustion waste gas of the reformer 10 and from the synthetic gas were supplied into the raw material gas introducing fluid passageway 20₂ under the conditions shown in Table 1 so as to carry out the steam reformation in the reaction tube 11 of the reformer 10, the crude methanol formation in the reaction apparatus 40 for synthesizing methanol, the recovery of the liquid crude methanol in the gas-liquid separator 61, and the recycle of the purge gas into the raw material gas line, thereby manufacturing methanol. Table 1 also shows the composition of the methanol thus manufactured.

[0043] Item (A) shown in Table 1 denotes the natural gas supplied to the raw material gas introducing fluid passageway 20₂. Item (B) represents the carbon dioxide gas recovered in the carbon dioxide recovery device 31 from the combustion waste gas generated from the combustion radiation section 12 of the reformer 10 and compressed by the compressor 32 so as to be supplied to the raw material gas introducing fluid passageway 20₂. Item (C) represents steam supplied to the raw material gas introducing fluid passageway 20₂. Item (D) represents the synthetic gas supplied to the heat recovery device 52 through the heat exchanger 51. Item (E) represents the synthetic gas supplied from the heat recovery device 52 into the compressor 53. Item (F) represents the unreacted gas (purge gas) separated in the gas-liquid separator 61. Item (G) represents a part of the purge gas separated in the gas-liquid separator 61 so as to be recycled into the raw material gas introducing fluid passageway 20₂. Further, item (H) shown in Table 1 represents the liquid crude methanol separated in the gas-liquid separator 61. Incidentally, these items (A) to (H) are shown in FIG. 1.

(Comparative Example 1)

[0044] Methanol was manufactured as in Example 1, except that used was a methanol manufacturing plant shown in FIG. 2. In this case, a carbon dioxide gas manufactured outside the plant was added to the natural gas through a fluid passageway 20₁₇, and the unreacted gas (purge gas) separated from the gas-liquid separator 61 was recycled to the fuel introducing fluid passageway 20₁ so as to be utilized as a fuel of the combustion radiation section 12 of the reformer 10. Table 2 shows the composition of the methanol thus obtained.

[0045] Item (A) shown in Table 2 represents the natural gas supplied into the raw material gas introducing fluid passageway 20₂. Item (B) represents the carbon dioxide gas supplied to the raw material gas introducing fluid passageway 20₂. Item (C) represents steam supplied into the raw material gas introducing fluid passageway 20₂. Item (D) represents the synthetic gas supplied to the heat recovery device 52 through the heat exchanger 51. Item (E) represents the synthetic gas supplied from the heat recovery device 52 into the compressor 53. Item (F) represents a part of the purge gas separated in the gas-liquid separator 61 so as to be recycled into the fuel introducing fluid passageway 20₁. Further, item (G) represents the liquid crude methanol separated in the gas-liquid separator 61. Incidentally, these items (A) to (G) are also shown in Table 2.

50

55

EP 1 182 185 A2

5	le 1)	(H)	9.0	J	0.3	J	0.4	20.1	78.6	100	1589
	(Example	(9)	12.3	ſ	84.2	1.3	1.6	ľ	0.6	100	2620
10		(F)	12.3	1	84.2	1.3	1.6	ı	0.6	100	2911
15		(E)	5.2	ı	75.1	14.0	5.2	0.5	-	100	7053
		(D)	3.9	_	56.4	10.5	3.9	25.3	t	100	9388
	Table 1	(C)	ı	-	-	1	-	100	ı	100	3446
25	H	(B)	1	1	1	ţ	100	1	ı	100	250
		(A)	94	9			1	r :	1	100	1000
30			CH4 (mol%)	C2H6 (mol%)	H2 (mol%)	CO (mol%)	CO2 (mo1%)	H ₂ O (mol%)	CH3OH (mol%)	Total (mol%)	:e 1r)
35		Item		<u> </u>			<u> </u>				Flow rate (kg·mol/hr)
40			Composition								ਸ਼ (

,										
le 1)	(<u>G</u>)	7.1	l	0.2	1	1.4	25.1	72.2	100	1566
Example	(F)	25.1	ı	66.8	1.7	5.8	ſ	9.0	100	422
ative	(E)	2.9		68.7	17.7	10.2	0.5	l	100	4268
(Comparative	(a)	1.9	ı	46.5	12.0	6.9	32.7	1	100	6305
2	(C)	1	ı		1	ı	100	ı	100	3180
Table	(B)	1	ı	ı	1	100	ı	ı	100	250
	(A)	94	9	,	- 2	,	,	ı	100	1000
		CH4 (mo1%)	C2H6 (mol%)	H2 (mol%)	CO (mol%)	CO ₂ (mol%)	H20 (mol%)	CH3OH (mol%)	Total (mol%)	te hr)
	Item	Composition								Flow rate (kg·mol/hr)

[0046] To reiterate, a part of the purge gas separated in the gas-liquid separator 61 is recycled in the method of the present invention (Example 1) into the raw material gas introducing fluid passageway 20₂ so as to be added to the natural gas having carbon dioxide and steam added thereto in advance. In Comparative Example 1, however, a part of the purge gas separated in the gas-liquid separator 61 is recycled as a fuel gas into the fuel gas introducing fluid passageway 20₁. As is apparent from Tables 1 and 2, Example 1 permits increasing the methanol production, compared with Comparative Example 1.

[0047] As described above, the present invention provides a method of manufacturing methanol, in which a purge gas mainly containing hydrogen, which is obtained by the gas-liquid separation from the crude methanol, is recycled so as to be added to the natural gas having steam and carbon dioxide added thereto in advance, thereby utilizing the purge gas as a part of the raw material. It follows that the methanol manufacturing method of the present invention permits increasing the methanol production.

Claims

1. A method of manufacturing methanol, characterized by comprising:

5

10

15

20

30

35

40

45

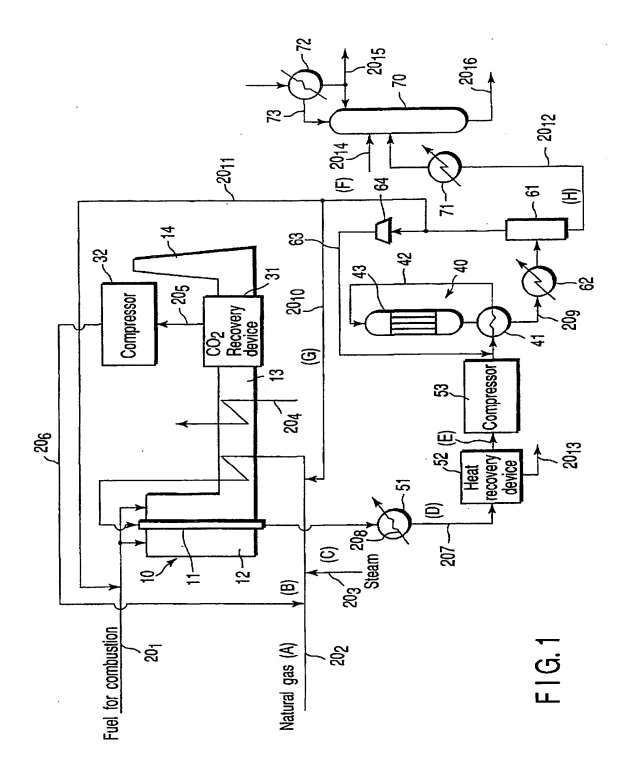
50

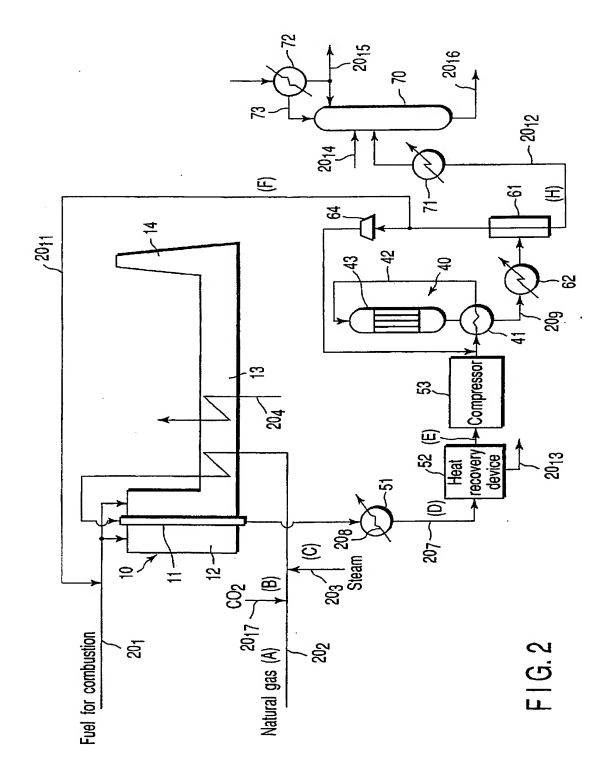
55

a synthetic gas forming step, in which a natural gas having steam and carbon dioxide added thereto is supplied into a reaction tube (11) of a reformer (10) provided with the reaction tube (11) and a combustion radiation section (12) for heating the reaction tube by the combustion of a fuel so as to bring about a reaction between the hydrocarbon contained in the natural gas and the steam, thereby forming a synthetic gas containing mainly hydrogen, carbon monoxide and carbon dioxide;

- a crude methanol synthesizing step for performing the reaction of the synthetic gas in the presence of a methanol synthetic catalyst so as to synthesize a crude methanol;
- a gas-liquid separation step for separating the crude methanol into a liquid crude methanol and a purge gas mainly containing hydrogen; and
- a distillation step for distilling the liquid crude methanol so as to separate a refined methanol; wherein a part of the purge gas is recycled into the natural gas having steam and carbon dioxide added thereto in advance.
- 2. The method of manufacturing methanol according to claim 1, characterized in that a part of all of the carbon dioxide gas added to the natural gas consists of a carbon dioxide gas recovered from at least one of the combustion waste gas generated from the combustion radiation section (12) of the reformer (10) and the combustion waste gas generated from a boiler for generating steam.
- 3. The method of manufacturing methanol according to claim 1, characterized in that a carbon dioxide gas and steam are added to the natural gas such that the molar ratio of methane (CH₄) contained in the natural gas to the steam (H₂O) falls within a range of between 1: 1.5 and 1: 5, and the molar ratio of methane (CH₄) contained in the natural gas to the carbon dioxide gas (CO₂) falls within a range of between 1: 1 and 1: 3.
- 4. The method of manufacturing methanol according to claim 1, characterized in that a part of the purge gas is recycled to the combustion radiation section (12) of the reformer (10) so as to be utilized as a part of the fuel.

9





Europäisches Patentamt

European Patent Office

Office européen des brevets



(11) EP 1 182 185 A3

(12)

EUROPEAN PATENT APPLICATION

(88) Date of publication A3: 13.08.2003 Bulletin 2003/33

(51) Int Cl.7: C07C 29/151, C01B 3/38

(43) Date of publication A2: 27.02.2002 Bulletin 2002/09

(21) Application number: 01119645.8

(22) Date of filing: 21.08.2001

(84) Designated Contracting States:

AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU

MC NL PT SE TR

Designated Extension States:

AL LT LV MK RO SI

(30) Priority: 21.08.2000 JP 2000249670

(71) Applicant: Mitsubishi Heavy Industries, Ltd. Tokyo (JP) (72) Inventors:

Kobayashi, Kazuto
 2-5-1, Marunouchi, Chiyoda-ku, Tokyo (JP)

lijima, Masaki
 2-5-1, Marunouchi, Chiyoda-ku, Tokyo (JP)

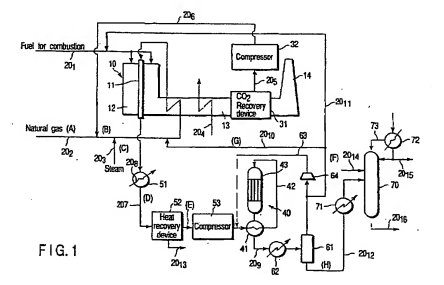
Morita, Kazuhiro
 2-5-1, Marunouchi, Chiyoda-ku, Tokyo (JP)

(74) Representative: HOFFMANN - EITLE Patent- und Rechtsanwälte Arabellastrasse 4 81925 München (DE)

(54) Method of manufacturing methanol

(57) A natural gas having steam and carbon dioxide added thereto is supplied into a reaction tube (11) of a reformer (10) provided with the reaction tube (11) and a combustion radiation section (12) for heating the reaction tube by combustion of a fuel so as to carry out a reaction between the hydrocarbon contained in the natural gas and the steam added to the natural gas, thereby forming a synthetic gas containing hydrogen, carbon

monoxide and carbon dioxide. Then, the synthetic gas is subjected to a reaction in the presence of a methanol synthesizing catalyst. A crude methanol obtained is subjected to a gas-liquid separation. Further, a liquid crude methanol separated in the separation is distilled. In the particular methanol manufacturing process, a part of the purge gas obtained in the separation is recycled into the natural gas having steam and carbon dioxide added thereto in advance.





EUROPEAN SEARCH REPORT

Application Number EP 01 11 9645

		ERED TO BE RELEVANT		
Category	Citation of document with inc of relevant passag		Helevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
Y	EP 1 008 577 A (MIT: 14 June 2000 (2000-0 * claim 1 *	SUBISHI HEAVY IND LTD 96-14)	1-4	C07C29/151 C01B3/38
Y	EP 0 650 950 A (MIT: CO) 3 May 1995 (1999 * claim 1 *	SUBISHI GAS CHEMICAL 5-05-03)	1-4	
				TECHNICAL FIELDS SEARCHED (Int.CL7)
				C01B C07C
	The present search report has I	been drawn up for all claims		
	Place of scarch	Date of completion of the search	1	Examiner
	MUNICH	27 June 2003	Ja	nus, S
X:par Y:par doo A;tec	ATEGORY OF CITED DOCUMENTS rticularly relevant if taken alone tricularly relevant if combined with anot sument of the same category shoological background n-written disclosure	E : earlier potent after the filing ber D : document oit L : document oit	ciple underlying the document, but pub- date ed in the application ed for other reasons to same patent tami	lished on, or

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 01 11 9645

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

27-06-2003

Patent document cited in search report		Publication date		Patent family member(s)	Publication date	
EP 1008577	A	14-96-2000	AU AU DE DK EP JP US JP	729993 B2 6437099 A 69905543 D1 1008577 T3 1008577 A1 2001097905 A 6218439 B1 2001097906 A	22-02-2001 22-06-2000 03-04-2003 26-05-2003 14-06-2000 10-04-2001 17-04-2001	
EP- 0650950	A	03-05-1995	JP AU CA DE DE EP US	7126201 A 680690 B2 7747694 A 2134406 A1 69407536 D1 69407536 T2 0650950 A1 5512599 A	16-05-1995 07-08-1997 18-05-1995 28-04-1995 05-02-1998 16-04-1998 03-05-1995 30-04-1996	

For more details about this annex : see Official Journal of the European Patent Office, No. 12/82